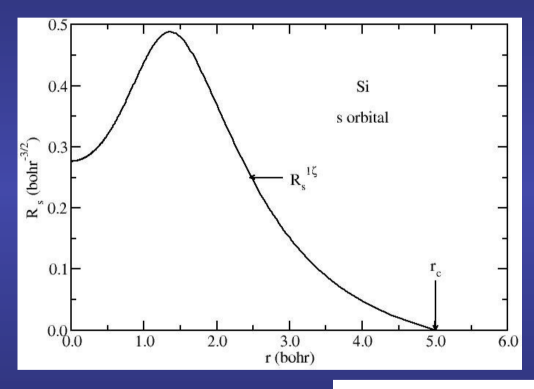
# Atomic orbitals of finite range as basis sets



Javier Junquera



### Most important reference followed in this lecture

phys. stat. sol. (b) 215, 809 (1999)

Subject classification: 71.15.Mb; 71.15.Fv; 71.24.+q; S1.3; S5; S5.11

### Linear-Scaling ab-initio Calculations for Large and Complex Systems

E. ARTACHO<sup>1</sup>) (a), D. SÁNCHEZ-PORTAL (b), P. ORDEJÓN (c), A. GARCÍA (d), and J. M. SOLER (e)

PHYSICAL REVIEW B, VOLUME 64, 235111

Numerical atomic orbitals for linear-scaling calculations

Javier Junquera, 1 Óscar Paz, 1 Daniel Sánchez-Portal, 2,3 and Emilio Artacho4

PHYSICAL REVIEW B 66, 205101 (2002)

Systematic generation of finite-range atomic basis sets for linear-scaling calculations

Eduardo Anglada, 1,2 José M. Soler, 1 Javier Junquera, 3 and Emilio Artacho 4

### ...in previous chapters:

### the many body problem reduced to a problem of independent particles

One particle Kohn-Sham equation

$$\left[ -\frac{1}{2} \nabla^2 + V_{eff}^{\sigma} \left( \vec{r} \right) \right] \psi_i^{\sigma} \left( \vec{r} \right) = \varepsilon_i^{\sigma} \psi_i^{\sigma} \left( \vec{r} \right)$$

$$V_{eff}^{\sigma}\left(\vec{r}\right) = V_{ext}\left(\vec{r}\right) + V_{Hartree}[n] + V_{xc}^{\sigma}[n^{\uparrow}, n^{\downarrow}]$$

Goal: solve the equation, that is, find

- the eigenvectors
- the eigenvalues

Solution: expand the eigenvectors in terms of functions of known properties (basis)

$$\psi_{i}\left(\vec{r}
ight)=\sum_{lpha}c_{ilpha}\left(\!f_{lpha}\left(\vec{r}
ight)\!
ight)$$

basis functions

### Different methods propose different basis functions

#### Each method has its own advantages:

- most appropriate for a range of problems
- provide insightful information in its realm of application

#### Each method has its own pitfalls:

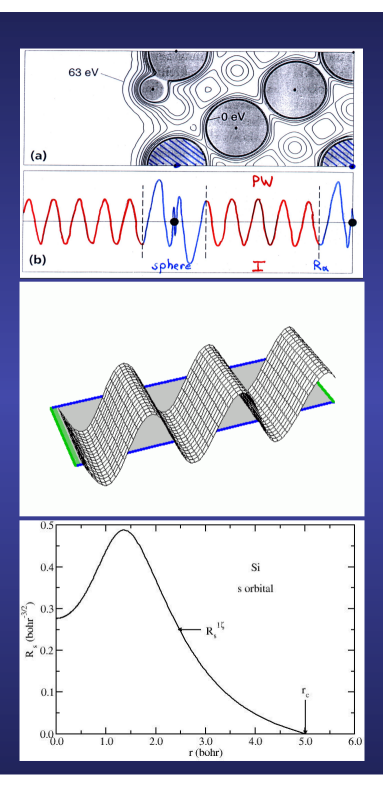
- importance to understand the method, the pros and the cons.
- what can be computed and what can not be computed

# Three main families of methods depending on the basis sets

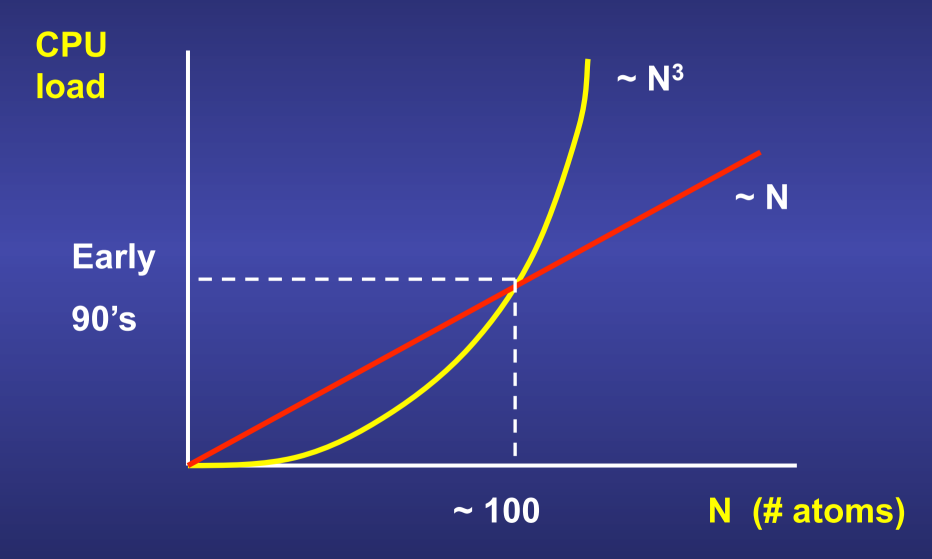
**Atomic sphere methods** 

Plane wave and grids

**Localized basis sets** 



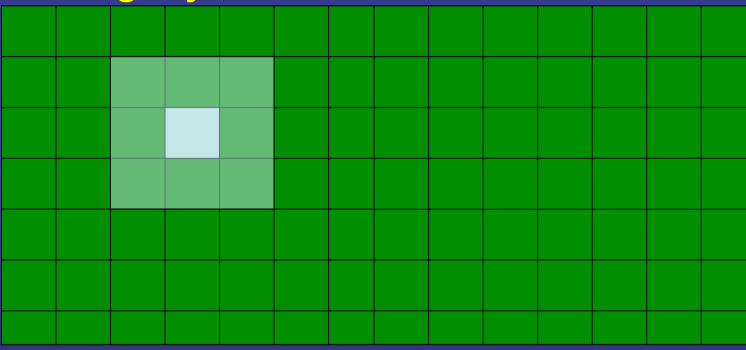
# Order-N methods: The computational load scales linearly with the system size



G. Galli and M. Parrinello, Phys. Rev Lett. 69, 3547 (1992)

# Locality is the key point to achieve linear scaling

#### Large system



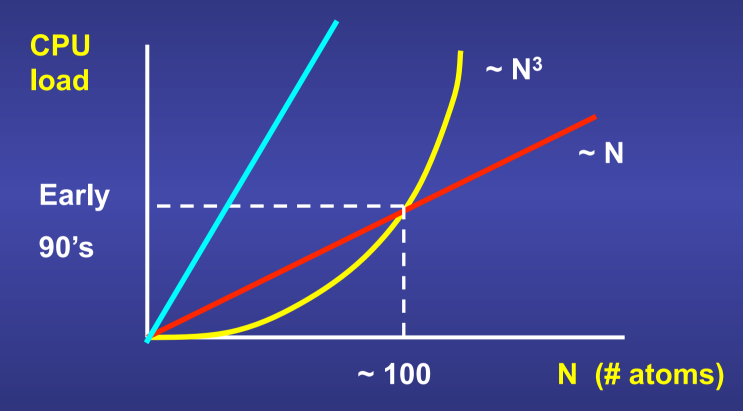
"Divide and Conquer"

W. Yang, Phys. Rev. Lett. 66, 1438 (1992)

**x2** 

# Efficient basis set for linear scaling calculations: localized, few and confined

**Locality** ⇒ Basis set of localized functions



Regarding efficiency, the important aspects are:

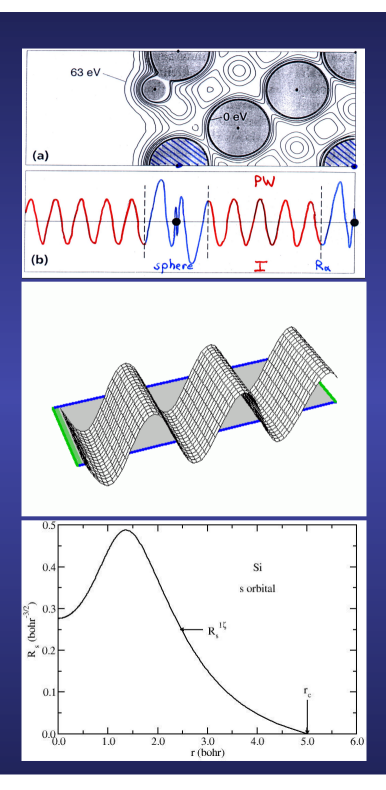
- NUMBER of basis functions per atom
- RANGE of localization of these functions

# Three main families of methods depending on the basis sets

**Atomic sphere methods** 

Plane wave and grids

**Localized basis sets** 



# **Atomic orbitals:** advantages and pitfalls

$$\phi_{Ilmn}(\vec{r}) = R_{Iln}(|\vec{r}_I|) Y_{lm}(\hat{r}_I)$$

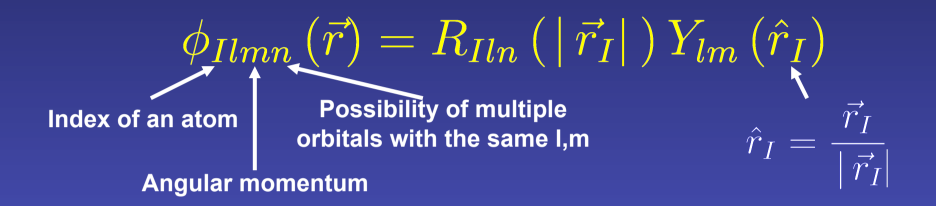
#### **ADVANTAGES**

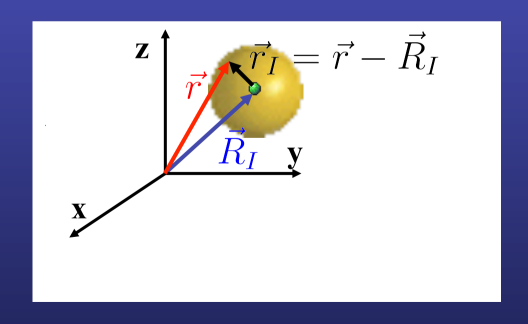
- Very efficient (number of basis functions needed is usually very small).
- Large reduction of CPU time and memory
- Straightforward physical interpretation (population analysis, projected density of states,...)
- They can achieve very high accuracies...

#### **DISADVANTAGES**

- ...Lack of systematic for convergence (not unique way of enlarge the basis set)
- Human and computational effort searching for a good basis set before facing a realistic project.
- Depend on the atomic position (Pulay terms).

# Atomic orbitals: a radial function times an spherical harmonic



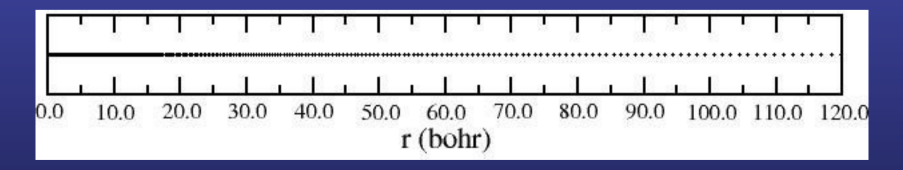


#### **Numerical atomic orbitals**

Numerical solution of the Kohn-Sham Hamiltonian for the isolated pseudoatom with the same approximations (xc,pseudos) as for the condensed system

$$\left(-\frac{1}{2r}\frac{d^2}{dr^2}r + \frac{l(l+1)}{2r^2} + V_l(r)\right)R_l(r) = \varepsilon_l R_l(r)$$

This equation is solved in a logarithmic grid using the Numerov method



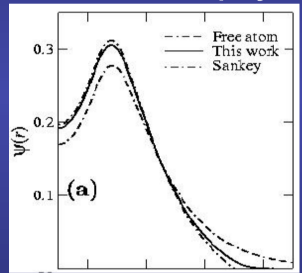
Dense close at the origin where atomic quantities oscillates wildly

Light far away from the origin where atomic quantities change smoothly

### Atomic orbitals: Main features that characterize the basis

$$\phi_{Ilmn}(\vec{r}) = R_{Iln}(|\vec{r}_I|) Y_{lm}(\hat{r}_I)$$

Radial part: degree of freedom to play with



Size: Number of atomic orbitals per atom

Range: Spatial extension of the orbitals

**Shape:** of the radial part

Spherical harmonics: well defined (fixed) objects



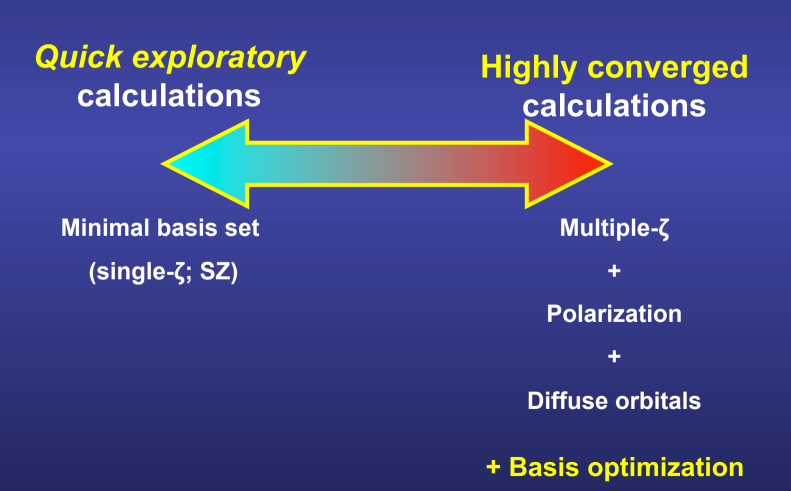






### Size (number of basis set per atom)

Depending on the required accuracy and available computational power



### Converging the basis size: from quick and dirty to highly converged calculations

Single-ζ (minimal or SZ)

One single radial function per angular

momentum shell occupied in the free-atom

#### Examples of minimal basis-set:

Si atomic configuration: 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup>

core

valence

$$l = 0 (s)$$

$$l = 1 (p)$$

$$m = 0$$

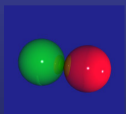
$$m = -1$$

$$m = 0$$

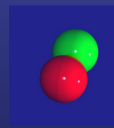
$$m = +1$$











4 atomic orbitals per Si atom

(pictures courtesy of Victor Luaña)

### Converging the basis size: from quick and dirty to highly converged calculations

Single-ζ (minimal or SZ)

One single radial function per angular

momentum shell occupied in the free-atom

#### **Examples of minimal basis-set:**

Fe atomic configuration: 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup> 3s2 3p6

4s<sup>2</sup> 3d<sup>6</sup>

core

valence

$$l=0$$
 (s)

m=0

$$m = -2$$

$$m = -1$$

$$m = 0$$

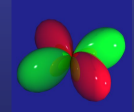
$$m = -2$$
  $m = -1$   $m = 0$   $m = +1$ 

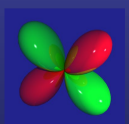
l = 2 (d)

$$m = +2$$











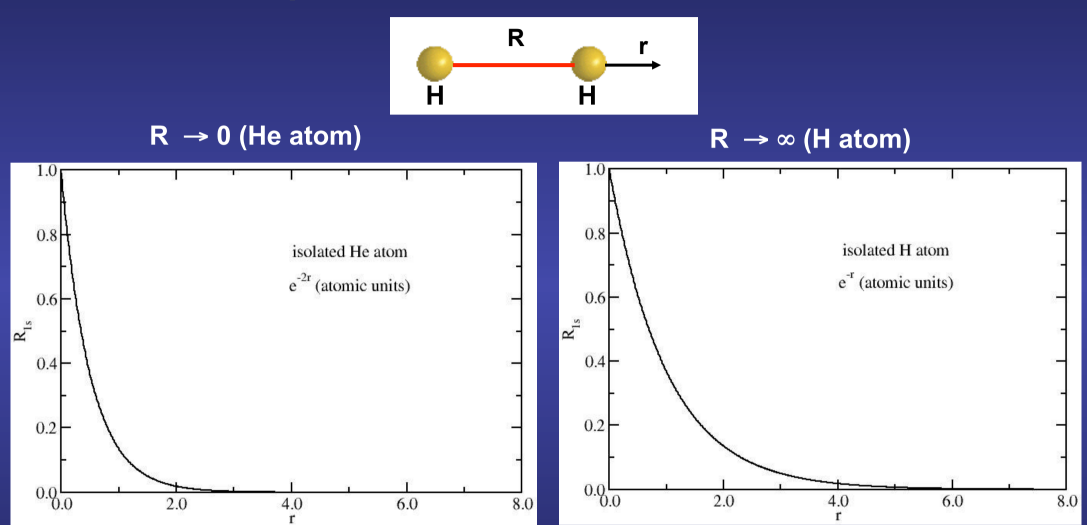




6 atomic orbitals per Fe atom

(pictures courtesy of Victor Luaña)

# The optimal atomic orbitals are environment dependent



Basis set generated for isolated atoms... ... but used in molecules or condensed systems

Add flexibility to the basis to adjust to different configurations

### Converging the basis size: from quick and dirty to highly converged calculations

Single-ζ (minimal or SZ)

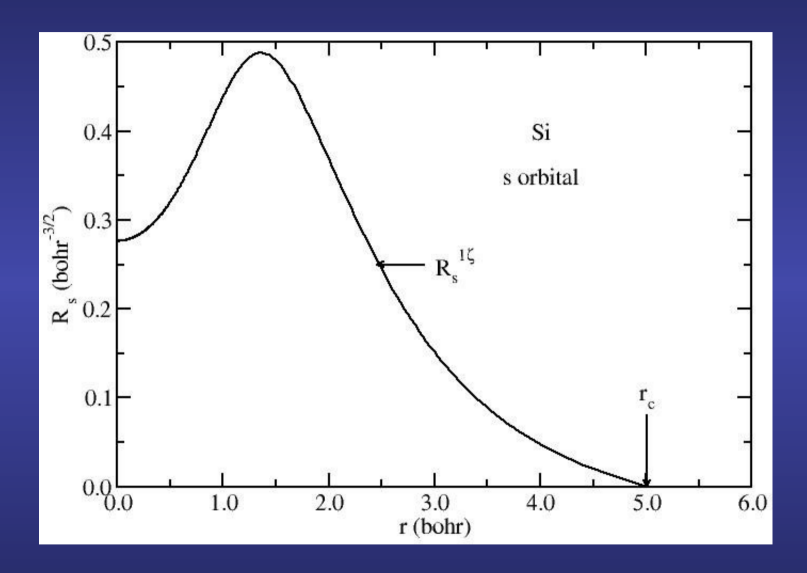
One single radial function per angular momentum shell occupied in the free-atom

Improving the quality

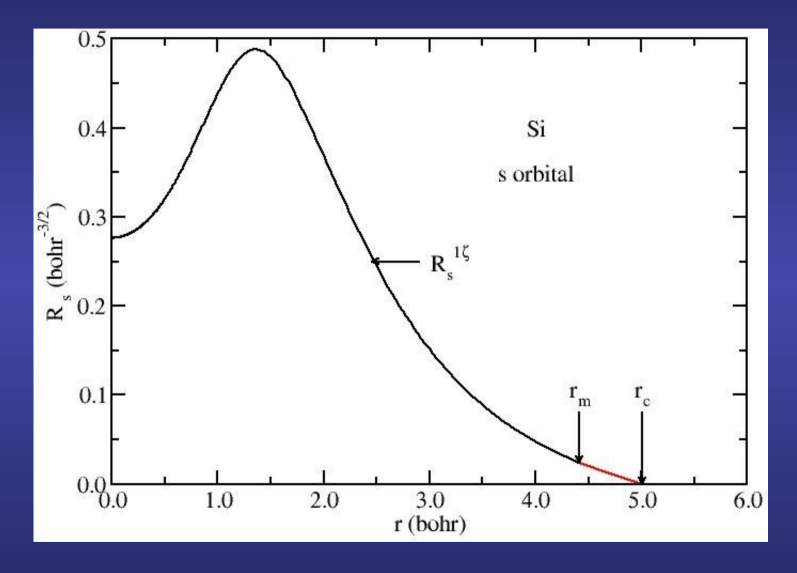
#### Radial flexibilization:

Add more than one radial function within the same angular momentum than SZ

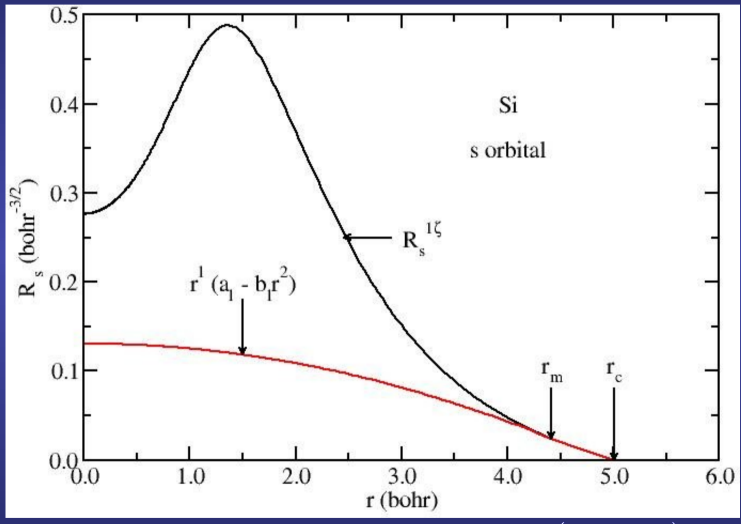
Multiple-ζ



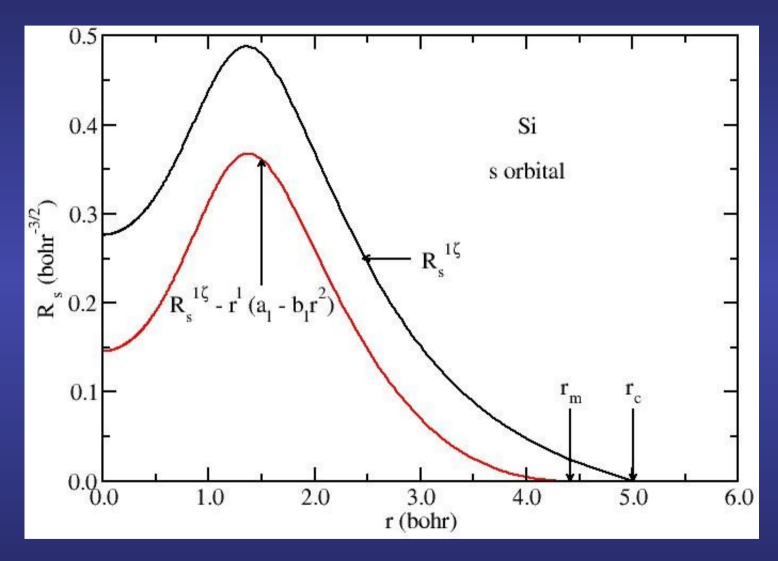
Starting from the function we want to suplement



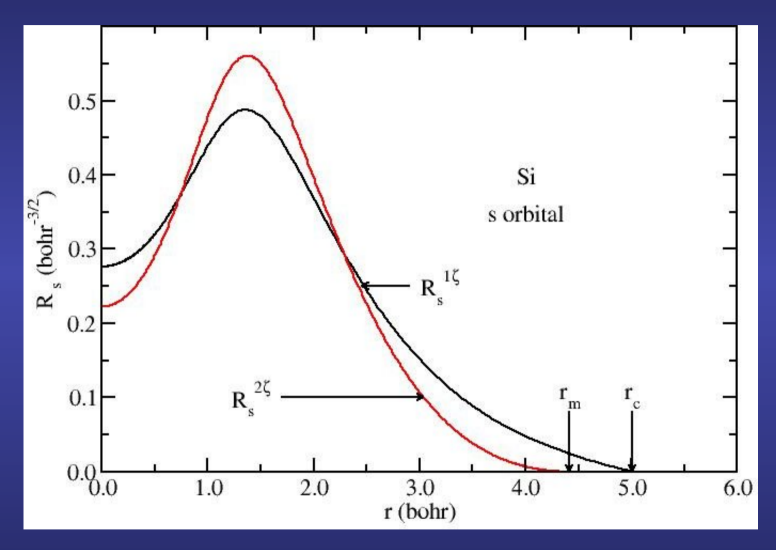
The second- $\!\zeta$  function reproduces the tail of the of the first- $\!\zeta$  outside a radius  $r_{m}$ 



And continuous smoothly towards the origin as  $r^l\left(a_l-b_lr^2
ight)$  (two parameters: the second- $\zeta$  and its first derivative continuous at  $r_{m}$ 



The same Hilbert space can be expanded if we use the difference, with the advantage that now the second- $\zeta$  vanishes at  $r_m$  (more efficient)



Finally, the second- $\zeta$  is normalized  $r_m$  controlled with PAO.SplitNorm (typical value 0.15)

### Converging the basis size: from quick and dirty to highly converged calculations

Single-ζ (minimal or SZ)

One single radial function per angular

momentum shell occupied in the free-atom

Improving the quality

#### Radial flexibilization:

Add more than one radial function within the same angular momentum than SZ

Multiple-ζ

#### **Angular flexibilization:**

Add shells of different atomic symmetry (different I)

**Polarization** 

### Example of adding angular flexibility to an atom Polarizing the Si basis set

Si atomic configuration: 1s<sup>2</sup> 2s<sup>2</sup> 2p<sup>6</sup>

core

$$l=0$$
 (s)

$$m = 0$$

m = -1

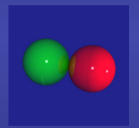
$$l=1$$
 (p)

$$m = 0$$

$$m = +1$$











#### Polarize: add l = 2 (d) shell

$$m = -2$$

$$m = -1$$

$$m = 0$$

$$m = -1$$
  $m = 0$   $m = +1$ 

$$m = +2$$









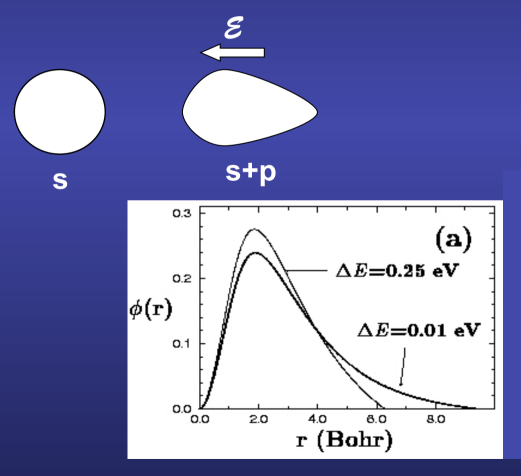


New orbitals directed in different directions with respect the original basis

# Two different ways of generate polarization orbitals

Perturbative polarization

Apply a small electric field to the orbital we want to polarize



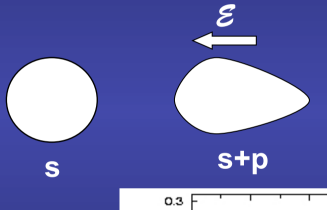
Si 3d orbitals

E. Artacho et al., Phys. Stat. Sol. (b), 215, 809 (1999)

# Two different ways of generate polarization orbitals

**Perturbative polarization** 

Apply a small electric field to the orbital we want to polarize



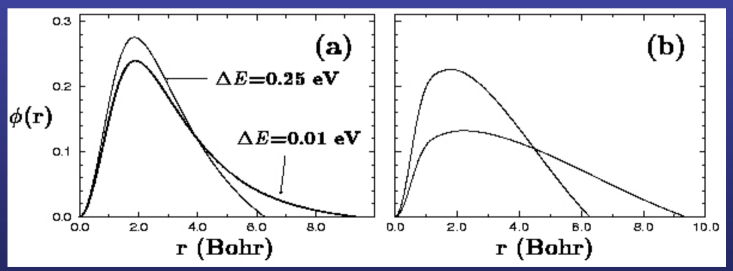
#### **Atomic polarization**

Solve Schrödinger equation for higher angular momentum

unbound in the free atom ⇒ require short cut offs

Si 3d

orbitals



E. Artacho et al., Phys. Stat. Sol. (b), 215, 809 (1999)

### Improving the quality of the basis ⇒ more atomic orbitals per atom

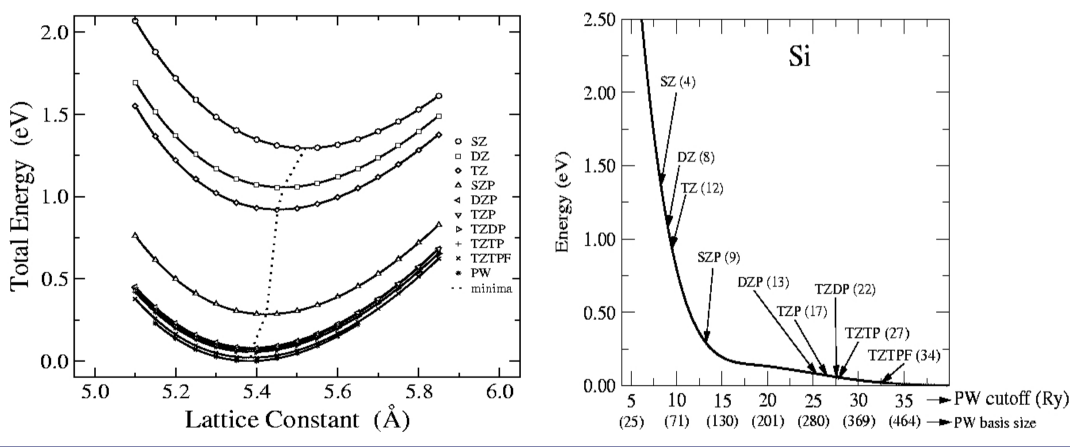
Atom	Valence	SZ			DZ	P		
	configuration							
		# orbita	ls symmetry	# orbital	ls symmetry	# orbitals	symmetry	
Si	$3s^2 \ 3p^2$	1	s	2	s	1	$d_{xy}$	
		1	$p_x$	2	$p_x$	1	$d_{yz}$	
		1	$p_y$	2	$p_y$	1	$egin{array}{c} d_{zx} \ d_{x^2-y^2} \ d_{3z^2-r^2} \end{array}$	
		1	$p_z$	2	$p_z$	1	$d_{x^2-y^2}$	
						1	$d_{3z^2-r^2}$	
	Total	4		8		(DZ+P) 13		

Atom	Valence						
	configuration						
		# orbita	ls symmetry	# orbitals	symmetry	# orbitals	symmetry
Fe	$4s^2 \ 3d^6$	1	s	2	$\boldsymbol{s}$	1	$p_x$
		1	$d_{xy}$	2	$d_{xy}$	1	$p_y$
		1	$d_{yz}$	2	$egin{aligned} d_{xy} \ d_{yz} \end{aligned}$	1	$p_{z}$
		1	$d_{zx}$	2	$d_{zx}$		
		1	$d_{zx} \ d_{x^2-y^2} \ d_{3z^2-r^2}$	2	$d_{zx} \ d_{x^2-y^2} \ d_{3z^2-r^2}$		
		1	$d_{3z^2-r^2}$	2	$d_{3z^2-r^2}$		
	Total	6		12		(DZ+P) 15	1

# Convergence as a function of the size of the basis set: Bulk Si

#### **Cohesion curves**

#### PW and NAO convergence



Atomic orbitals show nice convergence with respect the size

Polarization orbitals very important for convergence (more than multiple- $\zeta$ )

Double-ζ plus polarization equivalent to a PW basis set of 26 Ry

# Convergence as a function of the size of the basis set: Bulk Si

	SZ	DZ	TZ	SZP	DZP	TZP	TZDP	PW	APW	Exp
a (Å)	5.52	5.46	5.45	5.42	5.39	5.39	5.39	5.38	5.41	5.43
B (GPa)	89	96	98	98	97	97	96	96	96	98.8
E <sub>c</sub> (eV)	4.72	4.84	4.91	5.23	5.33	5.34	5.34	5.37	5.28	4.63

A DZP basis set introduces the same deviations as the ones due to the DFT or the pseudopotential approaches

SZ = single-ζ

**P=Polarized** 

PW: Converged Plane Waves (50 Ry)

**DZ=** doble- ζ

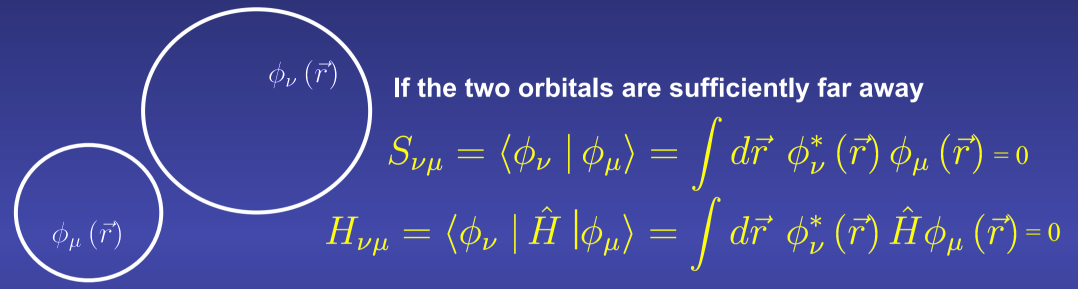
DP=Doble-polarized

**APW: Augmented Plane Waves** 

TZ=triple- ζ

# Range: the spatial extension of the atomic orbitals

Order(N) methods ⇒ locality, that is, a finite range for matrix and overlap matrices



#### **Neglect interactions:**

Below a tolerance

Beyond a given scope of neighbours

**Difficulty:** introduce numerical instabilities for high tolerances.

#### **Strictly localized atomic orbitals:**

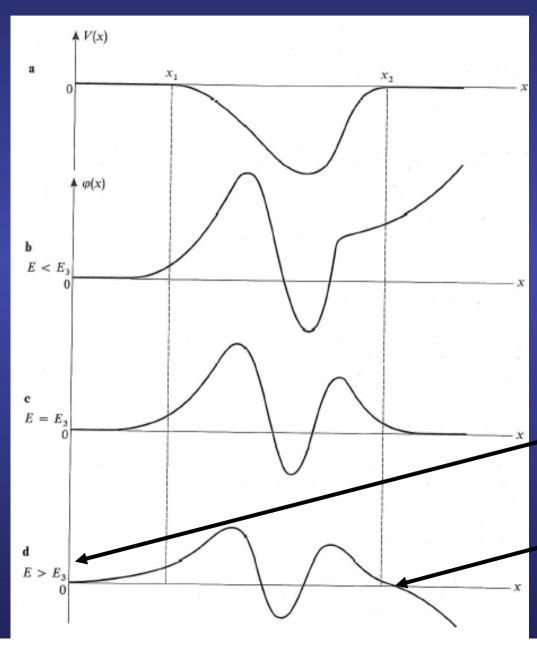
Vanishes beyond a given cutoff radius

O. Sankey and D. Niklewski, PRB 40, 3979 (89)

Difficulty: accuracy and computational efficiency depend on the range of the basis orbitals

How to define all the  $r_c$  in a balance way?

### How to control the range of the orbitals in a balanced way: the energy shift



### Particle in a confinement potential:

Imposing a finite 
$$\int_{-\infty}^{+\infty} |\phi(x)|^2 dx$$

Continuous function and first derivative



E is quantized (not all values allowed)

Increasing  $E \Rightarrow \phi_{\mu}$  has a node and tends to  $-\infty$  when  $x \rightarrow -\infty$ 

Complement M III "Quantum Mechanics", C. Cohen-Tannoudji *et al.* 

### How to control de range of the orbitals in a balanced way: the energy shift

$$\left(-\frac{1}{2r}\frac{d^2}{dr^2}r + \frac{l(l+1)}{2r^2} + V_l(r)\right)R_l(r) = (\varepsilon_l + \delta\varepsilon_l)R_l(r)$$

Energy increase ≡ Energy shift PAO.EnergyShift (energy)

Cutoff radius,  $r_c$ , = position where each orbital has the node

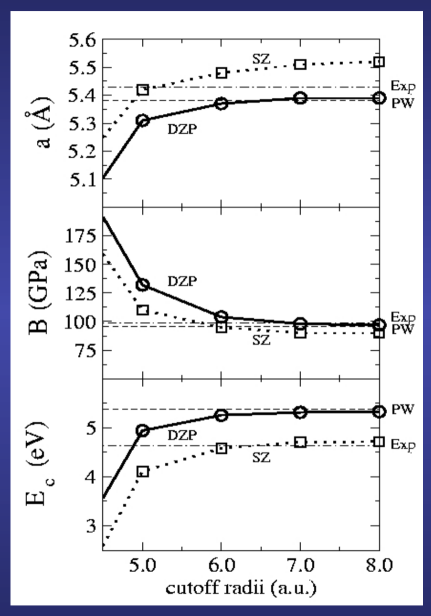
A single parameter for all cutoff radii

The larger the Energy shift, the shorter the r<sub>c</sub>'s

Typical values: 100-200 meV

E. Artacho et al. Phys. Stat. Solidi (b) 215, 809 (1999)

### Convergence with the range



**Bulk Si** 

equal *s*, *p* orbitals radii

J. Soler et al., J. Phys: Condens. Matter, 14, 2745 (2002)

More efficient

**More accurate** 

# The range and shape might be also controlled by an extra charge and/or by a confinement potential

#### Extra charge $\delta Q$

Orbitals in anions tend to be more delocalized

Orbitals in cations tend to be more localized

(For instance, this parameter might be important in some oxides)

#### **Confinement potentials**

Solve the Schrödinger equation for the isolated atom inside an confinement potential

Different proposals for the confinement potentials:

**Hard confinement** 

#### **Fireball**

O. F. Sankey and D. J. Niklewski, Phys. Rev. B 40, 3979 (89)

#### The default in SIESTA

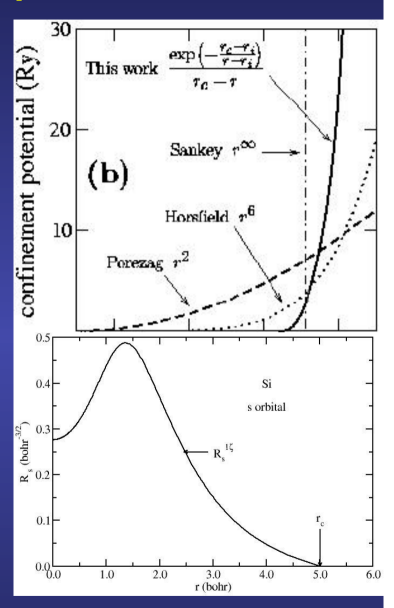
$$V = \infty, r \ge a$$

**Determined by the energy shift** 

$$V = 0, r < a$$

**Advantages:** empirically, it works very nice

Pitfall: produces orbitals with first derivative discontinuous at  $r_c$  problem when combined with numerical grids.

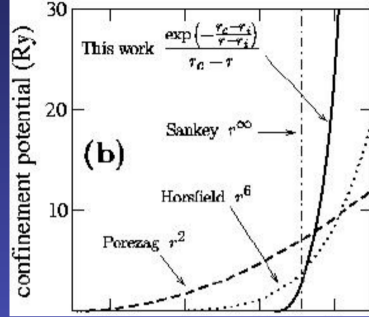


### Different proposals for the confinement potentials: Soft-confinement potential

#### **Available in SIESTA**

$$V\left(r\right) = V_0 \frac{e^{-\frac{r_c - r_i}{r - r_i}}}{r_c - r}$$

J. Junquera et al., Phys. Rev. B 64, 235111 (2001)



Advantages: orbital continuous with all the derivatives continuos

diverges at r<sub>c</sub> (orbital exactly vanishes there)

zero at the core region

Pitfall: two new parameters to play with, more exploratory calculations

### How to introduce the basis set in SIESTA Effort on defining a systematic with minimum parameters

If nothing is specified: default

#### **Default value**

Basis size: PAO.BasisSize DZP

Range of first-zeta: PAO.EnergyShift 0.02 Ry

Second-zeta: PAO.BasisType Split

Range of second-zeta: PAO.SplitNorm 0.15

**Confinement:** Hard well

Good basis set in terms of accuracy versus efficiency

### More global control on the basis with a few input variables: size and range

#### Size:

Basis size:

PAO.BasisSize SZ

DZ

SZP

**DZP** 

#### Range:

Range of first-zeta: PAO.EnergyShift 0.02 Ry

Range of second-zeta: PAO.SplitNorm 0.15

The larger both values, the more confined the basis functions

#### Some variable might be computed automatically

These variables calculated from PAO.EnergyShift and PAO.SplitNorm values

#### Adding polarization orbitals: perturbative polarization

```
%block PAO.Basis  # Define Basis set

H    1 +0.25  # Species label, number of 1-shells, charge
n=1    0    2 P    1  # n, l, Nzeta, Polarization, NzetaPol
    5.000    3.000  # rc (first-zeta), rm (second-zeta)
    1.000    1.000  # scaling factors
%endblock PAO.Basis
```

#### Adding polarization orbitals: atomic polarization

```
%block PAO.Basis
                            # Define Basis set
          +0.25
                            # Species label, number of 1-shells, charge
n=1 0 2
                            # n, l, Nzeta
                            # rc (first-zeta), rm (second-zeta)
  5.000
             3.000
                            # scaling factors
  1.000
             1.000
n=2 1 1
                            # n, l, Nzeta
                            # rc (first-zeta)
  5.000
                            # scaling factors
   1.000
%endblock PAO.Basis
```

#### **Soft-confinement potential**

```
%block PAO.Basis
                           # Define Basis set
 1 +0.25
                           # Species label, number of 1-shells, charge
n=1 0 2 E 150.00 4.5
                           # n, l, Nzeta, flag soft-conf, prefactor, inner rad
  5.000 3.000
                           # rc (first-zeta), rm (second-zeta)
  1.000 1.000
                           # scaling factors
%endblock PAO.Basis
                                  V_0 in Ry
```

 $r_i$  in bohrs

### Recap

#### **Numerical Atomic Orbitals**

A very efficient basis set

**Especially suitable for Order-N methods** 

Smooth transition from quick exploratory calculations to highly converged

Lack of systematic convergence

#### Simple handles for tuning the basis sets

Generate multiple-ζ: Split Valence

Generate polarization orbitals: Perturbative polarization

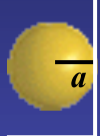
Control the range of the orbitals in a balanced way: Energy Shift

**Confine the orbitals: Soft-confinement potential** 

A DZP basis set, the same deviations as DFT functional or Pseudo

### **Suplementary information**

### Spherical Bessel functions $j_l(kr)$ , solutions of a free particle confined in a box



$$V = \infty, r \ge a$$

Schrödinger equation for a particle inside the box

$$-\frac{\hbar^2}{2m}\nabla^2\psi(r,\theta,\phi) = E\psi(r,\theta,\phi)$$

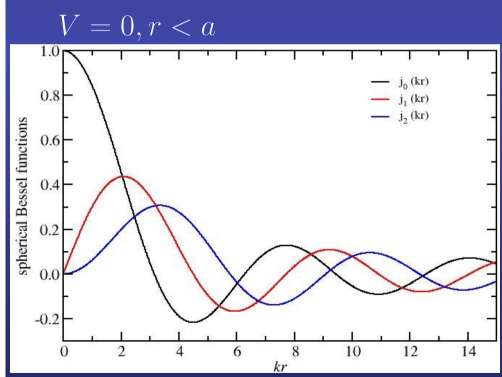
After separation of variables, the radial equation reads

$$\psi(r, \theta, \phi) = R(R)\Theta(\theta)\Phi(\phi)$$

$$\frac{d^2R}{dr^2} + \frac{2}{r}\frac{dR}{dr} + \left[k^2 - \frac{l(l+1)}{r^2}\right]R = 0$$
 
$$k^2 = \frac{2mE}{\hbar^2} \qquad \begin{array}{c} l \in \mathbf{Z}, \text{ separation} \\ \text{variable constant} \end{array}$$

Solution of the radial equation

$$R(r) = \begin{cases} Aj_l(kr) + Bn_l(kr), & r < a \\ 0, & r \ge a \end{cases}$$



Boundary conditions: k must satisfy  $j_l(ka) = 0$ 

Spherical von Neumann function, not finite at the origin